PATENT APPLICATION FILING ACKNOWLEDGMENT

08/475252

Mailing Date: June 7, 1995 / Express Mail #TP175007070	
File No.: 16355-002500	
Inventor(s): Julian Nikolchev and Dai Ton Title: CONTRACTOR MDB/gs	
Title: CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE OCCLUSION	
Combined Declaration & Power of Attorney XX Assignment	
XX Assignment XX Small Project	
No. Pages of Span Small Emity Decl.	
No. Pages of Claims:	1
So of Claims.	
No. Sheets Dwgs: 5 (informal)	
ABSTRACT PAGE:	
SSIGNMENT COVER SHEET. 2 (\$606/DA20-1430)	
SSIGNMENT COVER SHEET: 2 (\$40/DA20-1430) Please stamp the date of receipt of the control of the	
Please stamp the date of receipt of the enclosed documents and return this card addressee.	
addressee.	to

B1750	27358 US POST OFFICE TO ADDRESSEE	SEE REVERSE FOR SERVICE GUARANTEE INSURANCE COVERAGE AND CLAIMS. THANK YOU FOR CHOOSING EXPRESS
	ORIGIN Date In: 6 75 Postage Fostage Next Day Delivery Date In: 6 75 Time In: A.M. P.M. P.M. P.M. Weight Ibs. oz. International Country Code Next Day Delivery or Second Day Delivery Date In: 6 75 A.M. P.M. Return Receipt \$ C.O.D. J. Date Vol. Total Postage & Fees Military 2nd Day Or Military 3rd Day	Walver of Signature I wish delivery to be made without obtaining the signature of the addressee or the addressee's agent (if in the judgment of the delivery employee, the article can be left in a secure location) and I suthorize the delivery employee to sign that the shipment was delivered and understand that the signature of
For Customer Use	Express Mall Corporate Account No.: FROM: MDBarrish / File 16355-25 TOWNSEND & TOWNSEND HOUFCLE & CREW 379 LYTTON AVE 2ND FL HALO ALTO CA 94301-1431	and Indemnity (Domestic Only) TO: Telephone Number: HON COMMISSIONER OF FATERIES AND TRADEMARKS WASHINGTON DC 20231-0001

TOWNSEND and Townsend Steuart Street Tower One Market Plaza San Francisco, CA 9410		IE and CREW	-	Labe	No	TB 17502	7358 US
(41 <i>5</i>) 543-9600	·		Date of Dep	osit	June 7	. 1995	
PATENT APPLICATION COMMISSIONER OF Washington, D. C. 202 Sir: Transmitted herewith for	PATENT AND 7 31 r filing is the [X]	patent application,	States Posts Addressee" indicated abo	d Service service ove and i	e "Exp under : s addre	oress Mail F 37 CFR 1.10	with the United Post Office to 0 on the date omnissioner of C. 20231
[] continuation-in-part p Inventors: JULIAN NI			Ву	2	/es/	Gene	1
		RVICAL FALLOPI ICAL FALLOPIAN					
Enclosed are:	c) of [] formal !	[X] informal drawin	a(s)		REC	THIS NUMBER EIPT WILL BE MAIN	358 US
		Conceptus, Inc., a		ation	, ,	,, 55, 5	
		n & Power of Attorn					
[X] A [X] signed [] u	_						
[X] A Power of Attorn							
	•	nall entity status und	ler 37 CFR 1.9 ar	id 37 CF	R 1.27	•	1:4:
[] A certified copy of		lan 27 CED 1 07					application.
[] Information Disclos			ent application of t	he contin	uation-i	n-nart annlica	tion
[] Enclosed is a pentite	[] Enclosed is a petition to extend time to respond in the parent application of the continuation-in-part application.						
The filing fee has been o	calculated as show	n below:				OTHER TH	IAN A
	(Col. 1)	(Col. 2)	SMALL EN	TITÝ		SMALL EN	•
FOR:	NO. FILED	NO. EXTRA	RATE	FEE	OR	RATE	FEE
BASIC FEE				\$ 365	OR		\$730
TOTAL CLAIMS	35 -20=	* 15	15 x11=	\$ 165	OR	x22=	\$
INDEP CLAIMS	5 -3=	* 2	2 x38=	\$ 76	OR	x76=	\$
[] MULTIPLE DEPE	NDENT CLAIM	PRESENTED	+120=	\$ -	OR	+240=	\$
* If the difference in Col. 1 is less than zero, enter "0" in Col. 2			TOTAL	\$ 606	OR	TOTAL	\$
Please charge Deposit	Account No. 20-	1430 as follows:					
[X] Filing fee				\$	606.00	·	
[X] Any additional fees associated with this paper or during the pendency of this application							
[] The issue fee set in 37 CFR 1.18 at or before mailing of the Notice of Allowance, pursuant to 37 CFR 1.311(b).							
[] A check for \$ is enclosed.			•	Respectfully submitted,			
A duplicate of this shee	t is enclosed.		TOWNSEN	D and TC	WNSE	ND KHOURI	E and CREW
			rel	D C	Sail	<u> </u>	
Telephone:			Mark D. Ba	rrish			

(415) 543-9600 J:\16355\25APP.TRN

Reg. No. 36,443 Attorneys for Applicant

DECLARATION



My residence, post office address and citizenship are as stated below next to my name; I believe I am the original, first and sole inventor
(if only one name is listed below) or an original, first and joint inventor (if plural inventors are named below) of the subject matter which
is claimed and for which a patent is sought on the invention entitled: CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE
OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE ATTACHMENT the specification of which X is attached
hereto or was filed on as Application Serial No and was amended on (if applicable).

I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, Section 1.56. I claim foreign priority benefits under Title 35, United States Code, Section 119 of any foreign applications(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed.

Prior Foreign Application(s)

Country	Application No.	Date of Filing	Priority Claimed Under 35 USC 119
			Yes No
			Yes No

I claim the benefit under Title 35, United States Code, Section 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, section 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, section 1.56 which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

Application Serial No.	Date of Filing	Status
		Patented Pending Abandoned
		Patented Pending Abandoned

Full Name of Inventor 1	Last Name NIKOLCHEV	First Name JULIAN	Middle Name or I	nitial
Residence & Citizenship	City Portola Valley	State/Foreign Country California	Country of Citizer United States of	•
Post Office Address	Post Office Address 251 Durazno Way	City Portola Valley	State/Country California	Zip Code 94028
Full Name of Inventor 2	Last Name TON	First Name DAI	Middle Name or Initial	
Residence & Citizenship	City San Jose	State/Foreign Country California	Country of Citizen United States of	•
Post Office, Address	Post Office Address 1693 Flickinger Avenue	City San Jose	State/Country California	Zip Code 95131
Full Name of Inventor 3	Last Name	First Name	Middle Name or Initial	
Residence & Citizenship	City	State/Foreign Country	Country of Citizenship	
Post Office Address	Post Office Address	City	State/Country	Zip Code

I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Signature of Inventor 1	Signature of Inventor 2	Signature of Inventor 3
JULIAN NIKOLCHEV	DAI TON	<i>''</i>
Date 6/7/95	Date 6/7/95	Date

DEC.MRO 1/93 J:\16355\16355-25.DEC

POWER OF ATTORNEY BY ASSIGNEE

CONCEPTUS, INC. is the Assignee of	of the invention entitled: <u>CONTRACEPTIVE</u>		
TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVI	CES HAVING MECHANICAL FALLOPIAN TUBE		
ATTACHMENT, the specification of which X is attached her	reto or was filed on as Application		
Serial No.			
	•		
• • • •	Attorney has been reviewed by the undersigned. The		
undersigned certifies that to the best of the undersigned's knowled			
(whose title is supplied below) is empowered to act on behalf of	the Assignee.		
	ey(s) and/or agent(s) to prosecute this application and		
transact all business in the Patent and Trademark Office connect	ed therewith.		
James M. Heslin, Re	eg. No. 29 541		
Gary T. Aka, Reg. 1			
Robert C. Colwell, I			
Paul C. Haughey, Ro			
David N. Slone, Reg			
William M. Smith, H			
Mark D. Barrish, Re	-		
Send Correspondence to:	Direct Telephone Calls to:		
James M. Heslin, Esq.	(Name, reg. no., tele. no.)		
TOWNSEND and TOWNSEND KHOURIE and CREW			
One Market Plaza	James M. Heslin		
Steuart Street Tower, 20th Floor	Reg. No. 29,541		
San Francisco, CA 94105	(415) 326-2400		
·			

CONCEPTUS, INC.

(Signature)

Name: Julian Nikoleher

Title: VP

J:\16355\16355-25.PWR PWR.MRG 10/93

VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 CFR 1.9(f) & 1.27(e)) - SMALL BUSINESS CONCERN

	•		
Applicant or Pate Serial or Patent N		CHEV and DAI TON	
Filed or Issued:			
Title:		E TRANSCERVICAL FALLOPIAN TUBE OCC	
	DEVICES HAVIN	G MECHANICAL FALLOPIAN TUBE ATTAC	HMENT
I hereby declare t	hat I am		
	[] the owner of the	e small business concern identified below:	
	[X] an official of th	e small business concern empowered to act on be	chalf of the concern identified below:
NAME OF SMA	ALL BUSINESS CONCERN	CONCEPTUS, INC.	·
ADDRESS OF S	SMALL BUSINESS CONCE	RN 1021 Howard Avenue, San Carlos, Cali	fornia 95131
in 37 CFR 1.9(d) concern, including concern is the average of the pay periods	, for purposes of paying reduced the purposes of the affiliates, does not be the previous fiscal so of the fiscal year, and (2) co	uced fees to the United States Patent and Tradem not exceed 500 persons. For purposes of this stat year of the concern of the persons employed on a	ncern as defined in 13 CFR 121.12, and reproduced tark Office, in that the number of employees of the ement, (1) the number of employees of the business full-time, part-time or temporary basis during each irectly or indirectly, one concern controls or has the
to the invention,	entitled CONTRACEPTIV	•	small business concern identified above with regard CCLUSION DEVICES HAVING MECHANICAL described in
[X]	the specification filed here		
[]	application Serial No. Patent No.	, filed, issued	
L J	-	, Issued	
1.9(d), or a nonp *NOTE: Separate	rofit organization under 37 C	FR 1.9(e).	ot qualify as a small business concern under 37 CFR cation having rights to the invention averring to their
ADDRESS		· · · · · · · · · · · · · · · · · · ·	
NAME	[] INDIVIDUAL	[] SMALL BUSINESS CONCERN	[] NONPROFIT ORGANIZATION
ADDRESS			• •
	[] INDIVIDUAL	[] SMALL BUSINESS CONCERN	[] NONPROFIT ORGANIZATION
prior to paying, o		-	resulting in loss of entitlement to small entity status ue after the date on which status as a small entity is
to be true; and fu or imprisonment,	rther that these statements we or both, under section 1001 of	ere made with the knowledge that willful false sta	ements made on information and belief are believed tements and the like so made are punishable by fine h willful false statements may jeopardize the validity directed.
NAME OF PER	SON SIGNING SON IF OTHER THAN OW	NER JUDIAN Miko	ches
	PERSON SIGNING	1021 Howard Avenue, San Carlos, C	California 94070
SIGNATURE _	15/11	DATE	6/2/95
7		DATE	
1:1103511033-23.583			

FORM	PTO-	.1	5	9	5
FORM (Rev. 6	3-93)	•		_	_

RECORDATION FORM COVER SHEET PATENTS ONLY

U.S. DEPARTMENT OF COMMERCE Patent and Trademark Office

To the Honorable Commissioner of Patents and Trademarks.	Please record the attached original documents or copy thereof.	
1. Name of conveying party(ies):	2. Name and address of receiving party(ies):	
JULIAN NIKOLCHEV and DAI TON	Name: CONCEPTUS, INC.	
	Mairie.	
	Internal Address:	
Additional name(s) of conveying party(ies) attached? Yes X No		
	Charact Address 1021 Howard Avenue	
	Street Address: 1021 Howard Avenue	
3. Nature of conveyance:	·	
x Assignment Merger	City: San Carlos State: California ZIP: 94070	
Security Agreement Change of Name		
Other:		
Superdian Datas Israe 7, 1005		
Execution Date: June 7, 1995	Additional name(s) & address(es) attached? Yes X No	
	Additional name(s) & address(es) attached? Yes x No	
4. Application number(s) or patent number(s).		
If this document is being filed together with a new applican	tion, the execution date of the application is: June 7, 1995	
A. Patent Application No.(s)	B. Patent No.(s)	
Additional numbers atta	ached? Yes x No	
5. Name and address of party to whom correspondence concerning document should be mailed:	6. Total number of applications and patents involved: 1	
Name: James M. Heslin, Esq. TOWNSEND and TOWNSEND KHOURIE and CREW	7. Total fee (37 CFR 3.41): \$ 40.00	
Twentieth Floor Steuart Street Tower	Enclosed X Charge Fees to Deposit Account	
One Market Plaza San Francisco, California 94105-1492	Charge any additional fees associated with this paper or	
(415) 326-2400	during the pendency of this application, or credit any overpayment, to deposit account	
	0. Danis and a sure 20. 1420	
	8. Deposit account number: 20-1430	
	(Attach duplicate copy of this page if paying by deposit account)	
DO NOT U	SE THIS SPACE	
9. Statement and signature.		
To the best of my knowledge and belief, the foregoing inf	formation is true and correct and any attached copy is a true copy	
of the original document.		
Mark D. Barrish Name of Person Signing Signature	<u>June 7, 1995</u> Date	
	pages including cover sheet, attachments, and document: 3	
10. Change Correspondence Address to that of Part 57 x	Yes No	
OMB No. 0651-0011 (exp. 4/94)		
Do not detach this portion Mail documents to be recorded with required cover sheet information to:		
Commissioner of Box	Patents and Trademarks Assignments ton, D.C. 20231	
J:\16355\16355-25.ASN		

ASSIGNMENT OF PATENT APPLICATION

JOINT

WHEREAS, JULIAN NIKOLCHEV, 251 Durazno Way, Portola Valley, California 94028, and DAI TON, 1693 Flickinger Avenue, San Jose, California 95131, hereinafter referred to as "Assignors," are the inventors of the invention described and set forth in the below identified application for United States Letters Patent:

Title of the Invention:

CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE ATTACHMENT

Date(s) of execution of Declaration: _	June 7, 1995	
Filing date:	Serial No.:	; and

WHEREAS, CONCEPTUS, INC., a California corporation, located at 1021 Howard Avenue, San Carlos, California 94070, hereinafter referred to as "Assignee," is desirous of acquiring an interest in the invention and application and in any Letters Patent and Registrations which may be granted on the same;

For good and valuable consideration, receipt of which is hereby acknowledged by Assignors, Assignors have assigned, and by these presents do assign to Assignee all right, title and interest in and to the invention and application and to all foreign counterparts (including patent, utility model and industrial designs), and in and to any Letters Patent and Registrations which may hereafter be granted on the same in the United States and all countries throughout the world, and to claim the priority from the application as provided by the Paris Convention. The right, title and interest is to be held and enjoyed by Assignee and Assignee's successors and assigns as fully and exclusively as it would have been held and enjoyed by Assignors had this assignment not been made, for the full term of any Letters Patent and Registrations which may be granted thereon, or of any division, renewal, continuation in whole or in part, substitution, conversion, reissue, prolongation or extension thereof.

Assignors further agree that they will, without charge to Assignee, but at Assignee's expense, (a) cooperate with Assignee in the prosecution of U.S. Patent applications and foreign counterparts on the invention and any improvements, (b) execute, verify, acknowledge and deliver all such further papers, including patent applications and instruments of transfer and (c) perform such other acts as Assignee

lawfully may request to obtain or maintain Letters Patent and Registrations for the invention and improvements in any and all countries, and to vest title thereto in Assignee, or Assignee's successors and assigns.

IN TESTIMONY WHEREOF, Assignors have signed their names on the dates indicated.

Date: 6/7/95

TULIAN NIKOLCHEV

Date: 6/7/95

DAI TON

J:\16355\16355-25.ASN ASSIGN.MRG 5/94

PATENT APPLICATION

CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE ATTACHMENT

Inventors:

Julian Nikolchev, a citizen of The United States, residing at 251 Durazno Way, Portola Valley, California, 94028;

Dai Ton, a citizen of The United States, residing at 1693 Flickinger Avenue San Jose, California, 95131; and

Assignee:

CONCEPTUS, INC. 1021 Howard Avenue San Carlos, California 94070, a California corporation.

Status:

SMALL ENTITY

TOWNSEND and TOWNSEND KHOURIE and CREW Steuart Street Tower, 20th Floor One Market Plaza San Francisco, California 94105 (415) 326-2400

Attorney Docket No. 16355-25

CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE ATTACHMENT

BACKGROUND OF THE INVENTION

1. Field of the Invention

5

10

15

20

25

30

35

The present invention relates generally to contraception, and more particularly to intrafallopian contraceptive devices and nonsurgical methods for their delivery.

Worldwide demand exists for safe, effective methods of both contraception and permanent sterilization. Although a variety of contraception and sterilization methods are available, all of the existing methods have limitations and disadvantages. Thus, the need for additional safe, low cost, reliable methods of contraception and permanent sterilization, both in developed and less developed countries, is widely recognized.

Many presently available contraception methods require significant user involvement, and user non-compliance results in quite high rates of failure. While the theoretical effectiveness of existing contraceptives, including barrier methods and hormonal therapies, is well established, overcoming user noncompliance to improve overall efficacy has proven difficult.

One form of contraception which is less susceptible to user noncompliance is the intrauterine device (IUD). IUDs have been found to have higher rates of reliability, and are effective for a longer period of time, than most other commercially available contraceptives. Unfortunately, IUDs are also associated with serious infectious complications. For this reason, the use of IUDs within the United States has decreased dramatically. Additionally, IUDs are subject to unplanned expulsion, and must be removed due to excessive pain or bleeding in a percentage of cases, further reducing the

acceptance of the IUD as a contraceptive method. Interestingly, the efficacy of copper IUDs appears to be higher than that of non-metallic IUDs. The reason for this has not been fully explained.

Commercially available options for permanent sterilization include fallopian tube ligation and vasectomy. These methods are surgical, are difficult to reverse, and are not available to many people in the world. It is common knowledge that fertilization occurs in the fallopian tubes where the sperm and ovum meet. Tubal ligation avoids this by complete occlusion of the fallopian tubes.

It has previously been proposed to reversibly occlude the fallopian tubes, for example, by in vitro formation of an elastomeric plug, or otherwise anchoring a device on either side of the narrowest region of fallopian tube, called the "isthmus." Such fallopian tube occlusion methods appear promising; however, an unacceptably high percentage of the non-surgical devices proposed to date have become dislodged during previous studies. Even where non-surgical intrafallopian devices have remained in place, they have been found to be only moderately effective at preventing conception.

For these reasons, it would be desirable to provide effective, reliable intrafallopian devices for contraception and sterilization. It would be particularly desirable to provide highly effective intrafallopian devices which did not require surgery for placement. It would be especially desirable if such devices and methods allowed easy placement of the device, but were less susceptible to being dislodged than previously proposed non-surgical intrafallopian devices.

2. Description of the Related Art

5

10

15

20

30

35

The experimental use of a stainless steel intrafallopian device is described in *Transcatheter Tubal Sterilization in Rabbits*, Penny L. Ross, RT 29 "Investigative Radiology", pp. 570-573 (1994). The experimental use of an electrolytically pure copper wire as a surgical contraceptive intrafallopian device in rats was described in "Antifertility

Effect of an Intrafallopian Tubal Copper Device", D.N. Gupta, 14 Indian Journal of Experimental Biology, pp. 316-319 (May 1976).

U.K. Patent Application Pub. No. 2,211,095 describes a uterine screw plug for blocking the fallopian tube. European Patent Application Pub. No. 0,010,812 describes a device for placement in the oviducts having enlargements at either end for anchoring the device. The same device appears to be described in Netherlands Patent No. 7,810,696.

5

10

15

20

30

35

The use of tubal occlusion devices is described in "Hysteroscopic Oviduct Blocking With Formed-in-Place Silicone Rubber Plugs", Robert A. Erb, Ph.D., et al., The Journal of Reproductive Medicine, pp. 65-68 (August 1979). A formed-in-place elastomeric tubal occlusion device is described in U.S. Patent No. 3,805,767, issued to Erb. U.S. Patent No. 5,065,751, issued to Wolf, describes a method and apparatus for reversibly occluding a biological tube. U.S. Patent No. 4,612,924, issued to Cimber, describes an intrauterine contraceptive device which seals the mouths of the fallopian tubes.

German Patent No. 28 03 685, issued to Brundin, describes a device for plugging a body duct with a device which swells when in contact with a body fluid.

Alternative contraceptive devices are disclosed in copending U.S. Patent Application Serial No. _____ (attorney docket no. 16355-24), the full disclosure of which is herein incorporated by reference.

SUMMARY OF THE INVENTION

The present invention provides intrafallopian devices and methods for their placement to prevent conception. The intrafallopian devices of the present invention are transcevically delivered and mechanically anchored within the fallopian tube to provide long term contraception, or alternatively permanent sterilization, without the need for surgical procedures or the risks of increased bleeding, pain, and infection associated with intrauterine devices (IUDs).

The intrafallopian devices of the present invention generally comprise a structure having a lumen-traversing region with a helical outer surface. The helical surface is mechanically anchored by a resilient portion of the structure which is biased to form an enlarged secondary shape, preferably forming distal and proximal anchoring loops. The anchoring loops help prevent the helical outer surface from rotating out of position, and also directly deter axial motion within the fallopian tube.

5

10

15

20

30

35

The use of copper in the intrafallopian device of the present invention improves its efficacy as a contraceptive Devices formed from plastically deformable materials, method. however, are less readily restrained in the fallopian tube. Apparently, the large variation in the actual shape and dimensions of fallopian tubes does not provide reliable anchoring for a pre-formed deformable intrafallopian device. The intrafallopian device of the present invention therefore comprises a resilient structure, usually a metallic coil, which includes a copper alloy or plating, ideally comprising an alloy including at least 75% copper. The coil material typically includes beryllium, zinc, stainless steel, platinum, a shape memory alloy, such as Nitinol^M, or the like. Preferably, the coil is composed of an alloy of beryllium and copper. Although the present device will generally result in occlusion, it need not completely occlude the fallopian tube to prevent the meeting of the sperm and ovum. Instead, the presence of the copper on the resilient structure is sufficient to provide effective contraception.

Conveniently, the present invention further comprises non-surgical placement of such intrafallopian devices by transcervical introduction. The resilient structure is restrainable in a straight configuration, e.g., by use of a corewire, greatly facilitating and reducing the risks of introduction. Thus, the cost and dangers associated with existing surgical contraceptive and sterilization procedures are avoided.

In a first aspect, a contraceptive intrafallopian device according to the present invention comprises a proximal anchor, a distal anchor, and a lumen-traversing region extending between the anchors. The lumen traversing region has a helical outer surface and a cross-section which is smaller than the cross-sections of the proximal and distal anchors.

Preferably, the lumen-traversing region comprises a resilient structure, generally having a ribbon wound over the outer surface to form the helical shape. Anchoring is enhanced by a sharp outer edge on the ribbon. As described above, at least one of the proximal anchor, the distal anchor, and the lumen-traversing region preferably comprises copper. The proximal and distal anchors generally comprise a resilient structure biased to form an enlarged secondary shape, thereby allowing the device to be restrained in a straight configuration to facilitate transcervical introduction.

In another aspect, a contraceptive intrafallopian device according to the present invention comprises a primary coil having a proximal loop, a distal loop, and an intermediate straight section between the loops. A helical ribbon is wound over at least a portion of the intermediate section, forming a helical surface to mechanically anchor the device within the fallopian tube.

The ribbon of the present intrafallopian device generally protrudes sufficiently to firmly engage the tubal wall. Preferably, the ribbon has a width in the range between .005 and .1 inch, a thickness in the range between .001 and .2 inch, and a pitch in the range between .01 and .2 inch. The overall device geometry preferably facilitates introduction and retention, but is not large or rigid enough to interfere with internal tissue movements. Usually, the device has a length in the range between 1.5 cm and 7.5 cm when in a relaxed state, while the distal loop and the proximal loop have outer diameters of at least 3 mm. Preferably, the primary coil has an outer diameter in the range between .2 mm and 5 mm.

In another aspect, a system for delivering intrafallopian contraceptive devices according to the present invention comprises a primary coil having a proximal loop, a distal loop, and an intermediate straight section between the Additionally, a lumen extends from a proximal end of the proximal loop to near a distal end of the distal loop. A helical ribbon is wound over at least a portion of the intermediate section, forming a helical surface to mechanically anchor the device within the fallopian tube. corewire is removably disposed within the lumen of the primary The corewire restrains the primary coil in a straight configuration, facilitating trancervical introduction. Optionally, the corewire is threadably received by the primary coil. Alternatively, a release catheter is slidably disposed over the corewire proximally of the primary coil to restrain the primary coil while the corewire is withdrawn proximally from the fallopian tube.

5

10

15

20

25

30

35

The helical ribbon is anchored in the fallopian tube by the distal and proximal loops. The ribbon is set in the tubal wall while the device is restrained in a straight configuration over a corewire by torquing on the corewire. Withdrawing of the corewire then releases the anchors. The distal anchor is generally inserted into the ampulla, distal of the isthmus, while the proximal anchor is located in the ostium. These anchors prevent rotation of the device, and also help avoid axial movement.

In yet another aspect, an intrafallopian contraceptive method according to the principles of the present invention comprises restraining a resilient contraceptive structure in a straight configuration over a corewire, where the resilient structure includes a lumentraversing region having a helical outer surface. The resilient structure is transcervically introduced into a target region of a fallopian tube, typically in the region of the ostium, and the corewire is withdrawn from the resilient structure. The resilient structure is mechanically anchored within the fallopian tube, a portion of the resilient structure assuming an enlarged secondary shape which is larger

in cross-section than the fallopian tube. Optionally, an electric current is applied through the resilient structure to the fallopian tube, thereby effecting permanent sterilization.

BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 illustrates a first embodiment of a contraceptive intrafallopian device according to the present invention.

Fig. 2 illustrates a primary coil used in the contraceptive intrafallopian device of Fig. 1.

5

10

15

20

25

30

35

Fig. 3 illustrates a secondary coil which has been imposed on a primary coil as used in the contraceptive intrafallopian device of Fig. 1.

Fig. 4 illustrates a corewire for use with the contraceptive intrafallopian device of Fig. 1.

Fig. 5 is a cross-sectional view of a contraceptive delivery system having the contraceptive intrafallopian device of Fig. 1.

Fig. 6 illustrates an alternative embodiment of the present contraceptive intrafallopian device.

Fig. 7 illustrates a primary coil used in the contraceptive intrafallopian device of Fig. 6.

Fig. 8 schematically illustrates a contraceptive delivery system including the contraceptive intrafallopian device of Fig. 6.

Figs. 9 and 10 illustrates a method of delivery of a contraceptive intrafallopian device according to the present invention.

DETAILED DESCRIPTION OF THE SPECIFIC EMBODIMENT

The present invention encompasses a contraceptive intrafallopian device which can alternatively be used as both a permanent and a reversible means of contraception. The present contraceptive methods and devices minimize the danger of non-use which has limited the efficacy of prior art contraceptive techniques. Moreover, the location of the present devices within the fallopian tubes provides a reduced risk of the infectious complications, increased bleeding, and

pelvic pain associated with intrauterine devices (IUDs). The location and the novel shape of the present intrafallopian device provides significant advantages over IUDs, which have been found to be susceptible to unplanned expulsion and removal due to excessive pain and bleeding. The present invention takes advantage of the increase in effectiveness associated with copper IUDs, providing a resilient structure including copper which may be transcervically positioned without the need for surgery.

Although the present contraceptive method is included within a group of contraceptive techniques generally referred to as fallopian tube occlusion methods, the present invention does not necessarily rely solely on blocking the fallopian tube to prevent fertilization. Instead, contraception is apparently provided by disrupting of ovum transport, the process of fertilization, and/or cleavage of the ovum. While the effect that copper has on these processes is not fully understood, it does appear that copper intrafallopian devices offer potentially significant increases in effectiveness over intrafallopian devices formed of other materials. Optionally, the present invention further encompasses devices which promote the growth of tissue within the tube to induce tubal occlusion, further inhibiting conception.

Conveniently, the present resilient structures are adapted to be releasably affixed over a corewire, the corewire restraining the resilient structure in a straight configuration. As the resilient structure has an outer diameter when in the straight configuration which is less than the inner diameter of the fallopian tube, the catheter containing the present intrafallopian device is easily transcervically introduced.

The present invention is anchored within the isthmus of the fallopian tube, overcoming the unintended expulsion of the device and the resulting failure of the contraceptive method. Such intrafallopian device expulsion has been the single greatest factor limiting the efficacy of easily positioned intrafallopian contraceptive techniques. The

present intrafallopian devices are generally elongate resilient structures pre-formed into secondary shapes. These secondary shapes will preferably form anchors proximally and distally of the narrowest portion of the fallopian tube, called the isthmus. The secondary shape must have a larger outer diameter than the inner diameter of the isthmus.

5

10

15

20

30

35

The present device is generally readily removed by snaring the resilient structure near the proximal end and pulling proximally on the resilient structure, thereby straightening the resilient structure and allowing it to be withdrawn without injuring the fallopian tube. Alternatively, an electrical current is applied to the device after it is positioned within the fallopian tube, providing permanent sterilization.

Referring now to Fig. 1, a first embodiment of the present contraceptive intrafallopian device 10 is formed from a resilient primary coil 12. Primary coil 12 has a proximal end 14 and a distal end 16, the latter having an atraumatic endcap 18. Primary coil 12 further includes three portions: a proximal anchor portion 20, a distal anchor portion 22, and a lumen-traversing region 24. Proximal and distal anchors 20,22 are biased to form anchoring loops 26, as described hereinbelow.

Lumen-traversing region 24 comprises a substantially straight portion of primary coil 12. A ribbon 28 is wound over the outer surface of primary coil 12 to provide a helical shape. Ribbon 28 includes sharp outer edges 29, which firmly anchor lumen-traversing region 24 in the fallopian tube wall when torque is applied to intrafallopian device 10. The ribbon is preferably formed of a high strength biocompatible metal, ideally being stainless steel. The ribbon is attached to primary coil 12 at a proximal joint 30 and a distal joint 32, which may be formed of solder, heat-shrink tubing, or the like.

Referring now to Fig. 2, primary coil 12 is most easily formed in a straight configuration as a cylindrical coil or spring, preferably having an outer diameter in the range from .005 inch to .05 inch, and having a length in the

range from 20 mm to 150 mm. Ideally, primary coil 12 has an outer diameter in the range from .01 inch to .05 inch and a length in the range from 30 mm to 125 mm.

5

10

15

20

25

30

35

Preferably, primary coil 12 is formed from a beryllium copper alloy wire. Beryllium copper provides the resilience necessary to avoid expulsion of the device, and also provides the increased effectiveness of a copper contraceptive intrafallopian device. Such a beryllium copper wire will typically have a diameter from .002 inch to .01 inch. To provide the increased efficacy of a copper intrafallopian device, primary coil 12 preferably comprises an alloy including 75% copper. Alternatively, primary coil 12 is formed from a resilient metal, such as stainless steel, platinum, a shape memory alloy, or the like. If such materials are used, primary coil 12 is preferably plated with copper or a copper alloy or otherwise has copper attached.

Primary coil 12 includes a body winding 42 and a thread winding 44. Body winding 42 is formed with the minimum possible pitch to increase the stiffness of primary coil 12. Thread winding 44 will typically comprise from 0.1 cm to 2 cm adjacent to proximal end 14, and will have a pitch roughly twice that of body winding 42.

Referring now to Fig. 3, the proximal and distal anchors are formed by imposing a bent secondary shape on selected portions of primary coil 12. The secondary shape preferably comprises loops 26 formed by bending primary coil 12, and heat treating the primary coil while it is bent. A wide variety of secondary shapes may be used, including sinusoidal curves, alternating loops, or loops separated by straight sections so as to form a "flower coil," as more fully described in copending U.S. Patent Application Serial No.

, (Attorney Docket No. 16355-24) the full disclosure of which is herein incorporated by reference. In all cases, the bent secondary shape should have an outer cross-section 46 which is larger than the fallopian tube to provide effective anchoring.

Referring now to Fig. 4, a corewire 50 for use with intrafallopian device 10 (Fig. 1) comprises a resilient wire

52 which tapers towards a distal end 54. Wire 52 is sufficiently stiff to restrain intrafallopian device 10 in a straight configuration, typically comprising stainless steel, platinum, or the like. A short section of coil forms corewire threads 56 attached at threadjoint 58. Threads 56 match the windings and pitch of threadwindings 44 of primary coil 12.

5

10

15

20

30

35

Referring now to Fig. 5, an intrafallopian contraceptive system 60 comprises corewire 50 inserted within a lumen 62 through intrafallopian device 10. Intrafallopian device 10 is releasably attached by engaging thread windings 44 with threads 56. Thus, intrafallopian device 10 is disengaged by torquing a proximal end of corewire 50 once intrafallopian device 10 is in position.

Referring now to Fig. 6, an alternative embodiment of the present intrafallopian device is again formed from a resilient primary coil 112 having a proximal end 114 and a distal end 116. The former includes a friction fitting 115. Primary coil 112 again includes three portions: a proximal anchor portion 120, a distal anchor portion 122, and a lumentraversing region 124. Proximal and distal anchors 120, 122 are here biased to form opposed anchoring loops 26, thereby increasing the relaxed overall cross-section of the proximal and distal anchors. A ribbon 128 is wound over the outer surface of primary coil 112 to provide a helical shape, as described above.

Referring now to Fig. 7, primary coil 112 comprises a uniform body winding 142. The secondary shape is imposed on the straight cylindrical coil as opposed loops 126, or alternatively as multiple loops of a flower coil.

Referring now to Fig. 8, an intrafallopian contraceptive system using alternative intrafallopian device 100 includes a corewire 152 which tapers towards a distal end 154. Friction fitting 115 fittingly engages corewire 152, which restrains primary coil 112 in a straight configuration. A release catheter 164 is slidably disposed over corewire 152 proximally of alternative intrafallopian device 100, allowing the device to be released by withdrawing corewire 152 relative to the release catheter.

Use of the present contraceptive intrafallopian device will be described with reference to Figs. 9 and 10. A uterine introducer canula 70 is inserted transcervically through a uterus 72 to the region of an ostium 74. Alternatively, a hysteroscope may be used in place of canula 70.

5

10

15

20

Intrafallopian contraceptive system 60 is advanced distally of introducer cannula 70 and manuevered through the fallopian tube, preferably until intrafallopian device 10 extends distally of the isthmus. Optionally, intrafallopian contraceptive system 60 is self-guided, with corewire 52 bent near distal end 54 to assist intraluminal manuevering. Alternatively, a guide wire and catheter are advanced into the fallopian tube first, and the guide wire is replaced with intrafallopian contraceptive system 60. In either case, the intrafallopian device is axially positioned with lumentraversing region 24 within a target region 84 adjacent to isthmus 80. Preferably, at least one loop of distal anchor 22 is distal of target region 84, and at least one loop of proximal anchor 20 is proximal of target region 84 to form the distal and proximal anchor bends.

Once intrafallopian device 10 is properly positioned, corewire 50 is torqued to set ribbon 28 in the tubal wall. The corewire may then be unthreaded from intrafallopian device 10 by rotating the corewire in the opposite direction, disengaging threads 56 from thread windings 44. The corewire is then free to slide proximally, releasing the primary coil. As the distal end of the primary coil is released, a distal anchor bend 90 is formed. Similarly, a proximal loop forms a proximal anchor bend 92. 30 The anchor bends help to axially restrain the device within the fallopian tube, and also prevent rotation around the helical shape of lumen-traversing region 24. As seen in Fig. 10, the loops need not assume their relaxed form to provide effective distal or proximal anchors. 35

The present invention further encompasses permanent sterilization by passing a current through the corewire to the intrafallopian device prior to withdrawing the corewire.

Fallopian tube tissue in contact with the intrafallopian device is dessechated, and thus attached to the present intrafallopian device. This action also causes permanent tubal damage, leading to the formation of scar tissue which encapsulates the intrafallopian device and causes permanent occlusion of the tubal lumen. Clearly, the corewire/primary coil interface must be conductive to allow the present non-surgical method of permanent sterilization.

5

In conclusion, the present invention provides a

contraceptive intrafallopian device which may be positioned without surgery. While the above is a complete description of the preferred embodiments of the invention, various alternatives, modifications, and equivalents may be used. For example, a wide variety of secondary shapes, including open loops, continuous bends, sinusoidal curves, or the like, may be imposed on the primary coil. Therefore, the above description should not be taken as limiting the scope of the invention, which is defined instead solely by the appended claims.

WHAT IS CLAIMED IS:

9

and the distal cross-section.

- An intrafallopian contraceptive device 1. 1 comprising: 2 a proximal anchor having a proximal cross-section; 3 a distal anchor having a distal cross-section; and 4 a lumen-traversing region extending between the 5 proximal anchor and the distal anchor, the lumen traversing 6 region having a helical outer surface and a helical cross-7 section which is smaller than both the proximal cross-section 8
- 2. An intrafallopian contraceptive device as claimed in claim 1, wherein the lumen-traversing region comprises a resilient structure.
- 3. An intrafallopian contraceptive device as claimed in claim 2, wherein the lumen-traversing region further comprises a ribbon wound over the outer surface of the resilient structure.
- 4. An intrafallopian contraceptive device as claimed in claim 2, wherein the ribbon includes a sharp outer edge.
- 5. An intrafallopian contraceptive device as claimed in claim 1 wherein at least one of the proximal anchor, the distal anchor, and the lumen-traversing region comprises copper.
- 6. An intrafallopian contraceptive device as claimed in claim 1 wherein at least one of the proximal anchor and the distal anchor comprises a resilient structure biased to form a secondary shape.
- 7. An intrafallopian contraceptive device as claimed in claim 6, wherein the resilient structure comprises a primary coil.

- 8. An intrafallopian contraceptive device as claimed in claim 7, wherein the primary coil comprises a material selected from the group consisting of beryllium, stainless steel, platinum, and shape memory alloy.
- 9. An intrafallopian contraceptive device as claimed in claim 8, wherein the primary coil comprises an alloy including beryllium and copper.
- 10. An intrafallopian device as claimed in claim 7, wherein the primary coil comprises an alloy including at least 75% copper.
- 11. An intrafallopian contraceptive device as
 2 claimed in claim 1, wherein a lumen extends from a proximal
 3 end of the proximal anchor to near a distal end of the distal
 4 anchor.
- 13. An intrafallopian contraceptive device as claimed in claim 12, wherein the ribbon has a width in the

the intermediate section.

3

range between .005 and .1 inch.

- 14. An intrafallopian contraceptive device as claimed in claim 13, wherein the ribbon has a thickness in the range between .001 and .2 inch.
- 15. An intrafallopian contraceptive device as claimed in claim 12, wherein the ribbon has a pitch in the range between .01 and .2 inch.

- 16. An intrafallopian contraceptive device as claimed in claim 12, wherein the device has a length in the range between 1.5 cm and 7.5 cm when in a relaxed state.
- 17. An intrafallopian contraceptive device as claimed in claim 12, wherein the device comprises copper.
- 18. An intrafallopian contraceptive device as claimed in claim 17, wherein the primary coil comprises a material selected from the group consisting of beryllium, stainless steel, platinum, and shape memory alloy.
- 19. An intrafallopian contraceptive device as claimed in claim 18, wherein the primary coil comprises an alloy including beryllium and copper.
- 20. An intrafallopian contraceptive device as claimed in claim 12, wherein the primary coil includes a lumen which extends from a proximal end of the proximal loop to near the distal end of the distal loop.
- 21. An intrafallopian contraceptive device as claimed in claim 12, wherein the primary coil has an outer diameter in the range between .2 mm and 5 mm.
- 22. An intrafallopian contraceptive device as

 claimed in claim 12, wherein the distal loop and the proximal

 loop have outer diameters of at least 3 mm when in a relaxed

 state.
- 23. An intrafallopian contraceptive system
 2 comprising:

a primary coil having a distal loop, a proximal
loop, an intermediate straight section between the distal loop
and the proximal loop, and a lumen from a proximal end of the
proximal loop to near a distal end of the distal loop;
a helical ribbon wound over at least a portion of

a helical ribbon wound over at least a portion of

8 the intermediate section; and

a corewire removably disposed within the lumen of the primary coil, the corewire restraining the primary coil in a straight configuration.

- 24. An intrafallopian contraceptive system as claimed in claim 23, wherein the primary coil comprises copper.
- 25. An intrafallopian contraceptive system as claimed in claim 23, wherein the corewire is threadably received by the primary coil.
- 26. An intrafallopian contraceptive system as
 claimed in claim 23, further comprising a release catheter
 slidably disposed over the corewire proximally of the primary
 coil, the release catheter having a distal primary coil
 engaging surface for restraining the primary coil while the
 corewire is withdrawn proximally.
- 27. An intrafallopian contraceptive method comprising:

9

10

11

12

13

restraining a resilient structure in a straight
configuration over a corewire, the resilient structure
including a lumen-traversing region having a helical outer
surface;

transcervically introducing the resilient structure into a target region of a fallopian tube; and

withdrawing the corewire from the resilient structure to mechanically anchor the resilient structure within the fallopian tube, at least a portion of the resilient structure assuming a secondary shape which is larger in crosssection than the fallopian tube.

28. A method as claimed in claim 27, wherein the target region is adjacent to an ostium of the fallopian tube.

- 29. A method as claimed in claim 28, wherein the target region extends distally of an isthmus of the fallopian tube.
- 30. A method as claimed in claim 27, further comprising torquing the corewire to anchor the resilient structure, the helical shape having a sharp outer edge.
- 31. A method as claimed in claim 27, wherein the
 withdrawing step comprises forming a distal anchor from a
 portion of the resilient structure which is distal of the
 lumen-traversing region, and forming a proximal anchor from a
 portion of the resilient structure which is proximal of the
 lumen-traversing region, the distal portion and the proximal
 portion assuming the secondary shape.
- 32. A method as claimed in claim 27, wherein the withdrawing step comprises unthreading the corewire from the resilient structure.
- 33. A method as claimed in claim 27, wherein the withdrawing step comprises axially restraining the resilient structure with a release catheter, the release catheter being slidably disposed over the corewire proximally of the resilient structure.
- 34. A method as claimed in claim 27, further comprising applying an electrical current through the resilient structure to the fallopian tube to permanently prevent conception.

1	35. An intrafallopian sterilization method
2	comprising:
3	transcervically introducing a structure into a
4	target region of a fallopian tube, the structure being
5	releasably attached to a distal end of an elongate body;
6	applying an electrical current through the elongate
7	body to the structure, and through the structure to the
8	fallopian tube to permanently anchor the structure within the
9	fallopian tube; and
10	releasing the structure from the elongate body and
11	withdrawing the elongate body.

CONTRACEPTIVE TRANSCERVICAL FALLOPIAN TUBE OCCLUSION DEVICES HAVING MECHANICAL FALLOPIAN TUBE ATTACHMENT

5

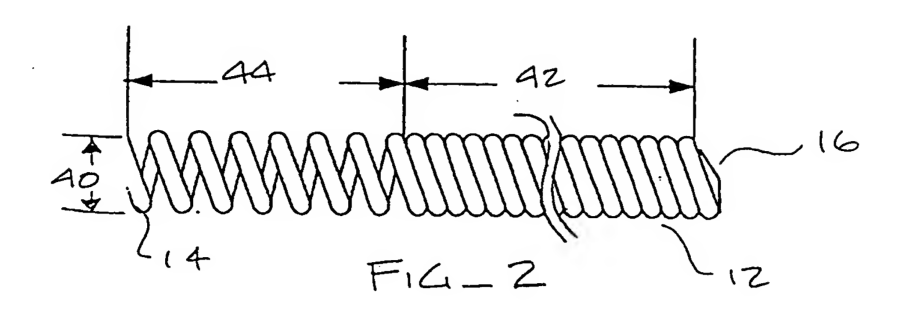
10

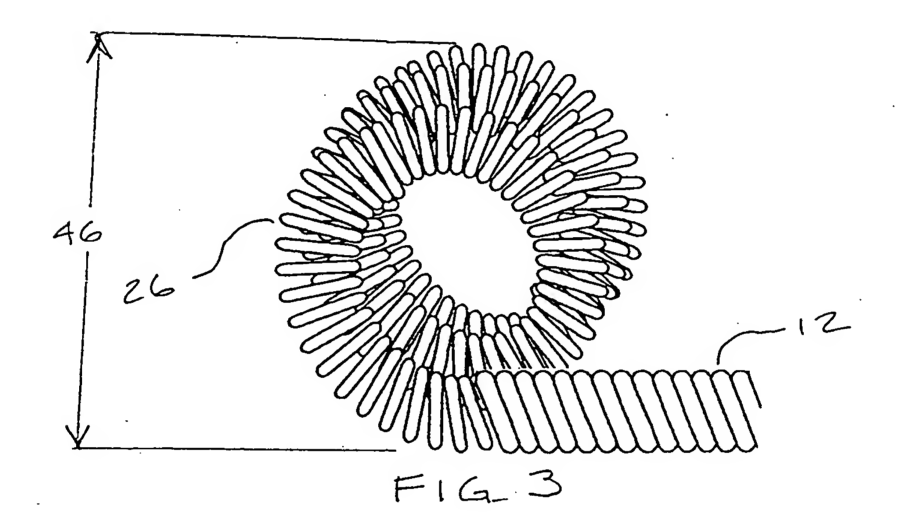
15

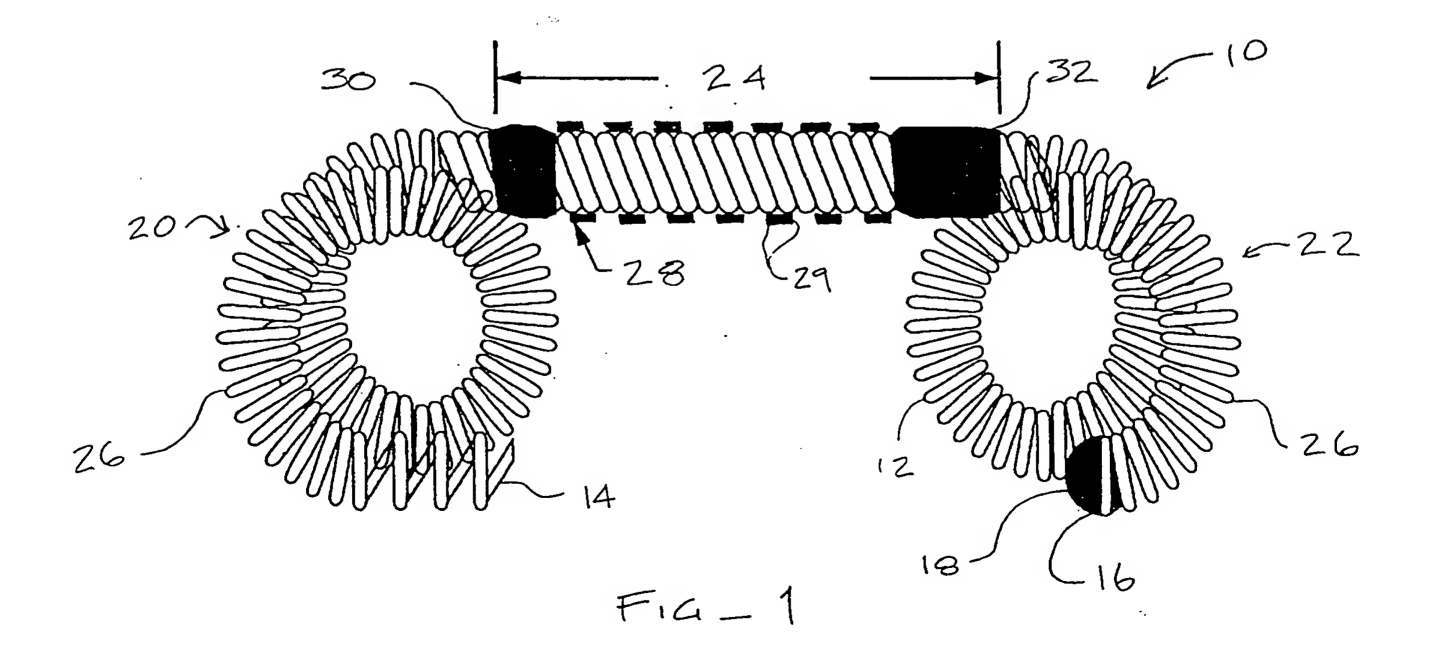
ABSTRACT OF THE DISCLOSURE

non-surgical methods for their placement to prevent conception. The efficacy of the device is enhanced by forming the structure at least in part from copper or a copper alloy. The device is anchored within the fallopian tube by a lumentraversing region of the resilient structure which has a helical outer surface, together with a portion of the resilient structure which is biased to form a bent secondary shape, the secondary shape having a larger cross-section than the fallopian tube. The resilient structure is restrained in a straight configuration and transcervically inserted within the fallopian tube, where it is released. Optionally, permanent sterilization is effected by passing a current through the resilient structure to the tubal walls.

20







 $\frac{F_{14}-7}{112}$

